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A

FISH & RICHARDSON P.C.

Frederick P. Fish
1855-1930

W.K. Richardson
1859-1951

February 18, 2000

225 Franklin Street
Boston, Massachusetts
02110-2804

Telephone
617 542-5070

Facsimile
617 542-8906

Web Site
www.fr.com

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Box Patent Application

Assistant Commissioner for Patents
Washington, DC 20231

Presented for filing is a new original patent application of:

Applicant: HUGH S. KEEPING and JONATHAN S. REICHNER

Title: TREATMENT FOR BONE DISORDERS

Enclosed are the following papers, including those required to receive a filing date under 37 CFR 1.53(b):

	<u>Pages</u>
Specification	21
Claims	4
Abstract	1
Declaration	[To be Filed at a Later Date]
Drawing(s)	5

Enclosures:

— Postcard.

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This application is entitled to small entity status. A small entity statement will be filed at a later date.

Basic filing fee	\$345
Total claims in excess of 20 times \$9	\$162
Independent claims in excess of 3 times \$39	\$78
Fee for multiple dependent claims	\$0
Total filing fee:	\$585

A check for the filing fee is enclosed. Please apply any other required fees or any credits to deposit account 06-1050, referencing the attorney docket number shown above.

If this application is found to be incomplete, or if a telephone conference would otherwise be helpful, please call the undersigned at (617) 542-5070.

Kindly acknowledge receipt of this application by returning the enclosed postcard.

Please send all correspondence to:

INGRID A. BEATTIE, PH.D., J.D.
Fish & Richardson P.C.
225 Franklin Street
Boston, MA 02110-2804

Respectfully submitted,



Ingrid A. Beattie, Ph.D., J.D.
Reg. No. 42,306
Enclosures
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APPLICATION
FOR
UNITED STATES LETTERS PATENT

TITLE: TREATMENT FOR BONE DISORDERS

APPLICANT: HUGH S. KEEPING AND JONATHAN S. REICHNER

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TREATMENT FOR BONE DISORDERS

BACKGROUND OF THE INVENTION

The invention relates to bone loss disorders. Catastrophic bone loss, i.e., osteolysis,
5 is a debilitating pathological consequence of a spectrum of disease states including
rheumatoid arthritis, osseous metastasis, aseptic prosthetic loosening and periodontitis.
Rheumatoid arthritis (RA) is a chronic inflammatory disease which often results in long term
disability and increased mortality.

SUMMARY OF THE INVENTION

10 The invention provides compositions and methods to deliver an anti-inflammatory
composition, e.g., recombinant human interleukin-4 (rhIL-4), to build (or rebuild) bone
tissue. The composition is produced from living osteoprogenitor cells (OPCs) or
odontoprogenitor cells. The cells contain a genetically-engineered viral or non-viral plasmid
vector containing a regulatable, inducible, osteoblast-specific promoter to direct expression
15 of an anti-inflammatory polypeptide at specific sites of implantation in bone to inhibit
osteolysis. For example, a bone stromal cell is isolated from autologous or allogeneic
periodontal ligament and manipulated *ex vivo* prior to implantation into a recipient patient.
Stromal cells are cultured in the presence of extracellular matrix (ECM) components to
differentiate into odontoprogenitor cells. For example, ECM contains bone morphogenic
20 proteins (BMPs) such as BMP-6. Induction of differentiation to progenitor cells is carried
out before or after genetic manipulation of the cells.

Preferably, the nucleic acid with which the cells are transfected or transduced encodes
an anti-inflammatory cytokine or anti-inflammatory fragment of the cytokine. For example,
the cytokine is interleukin-4 (IL-4). The nucleic acid encodes a polypeptide containing the
25 amino acid sequence of SEQ ID NO:1; for example, the nucleic acid contains the coding
region of the nucleotide sequence of SEQ ID NO:2.

Table 1: Human IL-4 Amino Acid Sequence

MGLTSQLLPPPLFFLLACAGNFVGHKCDITLQEIIKTLNSLTEQKTLCTELTVTDIFAA
 SKNTTEKEFCRAATVLRQFYSHHEKDTRCLGATAQQFHRHKQLIRFLKRLDRNLW
 GLAGLNSCPVKLEANQSTLENFLERLKTMREKYSKCSS (SEQ ID NO:2, GENBANK™
 5 Accession No. M13982)

Table 2: Human IL-4 Nucleotide Sequence

1 gatcgtagc ttctccgtat aaactaatttgc cctcacatttgc tcactgcaaa tcgacaccta
 61 ttaatgggttc tcacccccc actgcgtttcc cctctgttct tcctgttagc atgtccggc
 10 121 aactttgtcc acggacacaa gtgcgatatc accttacagg agatcatcaa aactttgaac
 181 agcctcacag agcagaagac tctgtgcacc gagttgaccg taacagacat ctttgctgcc
 241 tccaagaaca caactgagaaa ggaaaccttc tgcaaggctg cgactgtgct ccggcagttc
 301 tacagccacc atgagaagga cactcgctgc ctgggtgcga ctgcacagca gtcccacagg
 361 cacaaggcgc tgatccgatt cctgaaacgg ctgcacagga acctctgggg cctggcgggc
 421 ttgaatttctt gtcctgtgaa ggaagccaaac cagagtacgt tggaaaactt cttggaaagg
 481 ctaaagacga tcatgagaga gaaatattca aagtgtcgat gctgaatattt ttaatttatg
 541 agttttgtat agctttatattt ttaagtattt tatatatatttta taactcatca taaaataaaag
 601 tatatataga atct SEQ ID NO:2, GENBANK™ Accession No. M13982; coding

sequences span nucleotides 64-525; signal peptide encoded by nucleotides 64-135).

20 Alternatively, the cells contain a nucleic acid encoding an IL-4 fragment, agonist or mutant. The fragment, agonist or mutant has anti-inflammatory activity. For example, the mutant contains a mutation in the region of IL-4 which is involved in binding to IL-2R gamma, e.g., Arg 21 is changed to a Glu residue. Sequences which differ from the coding sequence of SEQ ID NO:2 hybridize under stringent conditions, with all or part of the reference sequence and encode an anti-inflammatory polypeptide. Promoter or transcriptional regulatory elements which differ from a reference sequence hybridize under stringent conditions to a nucleic acid having the reference sequence and retain transcription regulatory function, e.g., cell specificity, of the reference sequence. For example, the nucleic acid may contain one or more sequence modifications in relation to a reference sequence. Such modifications may be obtained by mutation, deletion and/or addition of one or more nucleotides compared to the reference sequence. Modifications are introduced to alter the activity of the regulatory sequence, e.g., to improve promoter activity, to suppress a transcription inhibiting region, to make a promoter constitutive or regulatable or vice versa.

25 30 35 Modifications are also made to introduce a restriction site to facilitate subsequent cloning

steps, or to eliminate the sequences which are not essential to the transcriptional activity. Preferably, a modified sequence is at least 70% (more preferably at least 80%, more preferably at least 90%, more preferably at least 95%, more preferably at least 99%) identical to a reference sequence. The modifications do not substantially alter the biological function 5 of a polypeptide or the cell-specificity of transcription promoter function associated with the reference sequence.

Nucleotide and amino acid comparisons are carried out using the Lasergene software package (DNASTAR, Inc., Madison, WI). The MegAlign module used was the Clustal V method (Higgins et al., 1989, CABIOS 5(2):151-153). The parameters used were gap 10 penalty 10, gap length penalty 10.

Alternatively, nucleic acids which differ from a given reference sequence hybridize at high stringency to a strand of DNA having the reference sequence, or the complement thereof. Hybridization is carried out using standard techniques, such as those described in Ausubel et al. (Current Protocols in Molecular Biology, John Wiley & Sons, 1989). "High 15 stringency" refers to nucleic acid hybridization and wash conditions characterized by high temperature and low salt concentration, i.e., hybridization at 42 degrees C, and in 50% formamide; a first wash at 65 degrees C, 2 × SSC, and 1% SDS; followed by a second wash at 65 degrees C and 0.2% × SSC, 0.190 SDS. Lower stringency conditions suitable for detecting DNA sequences having about 50% sequence identity to a reference gene or 20 sequence are detected by hybridization at 42 degrees C in the absence of formamide; a first wash at 42 degrees C in 6 × SSC and 1% SDS; and a second wash at 50 degrees C in 6 × SSC and 1% SDS.

A heterologous nucleic acid encoding a polypeptide (e.g., IL-4) is operably linked to an osteoblast-specific promoter such as an osteocalcin promoter sequence (e.g., a nucleic 25 acid containing the nucleotide sequence of SEQ ID NO:3) or a bone sialoprotein promoter sequence (nucleotides 1-2472 of SEQ ID NO:4) or dentin sialoprotein promoter sequence (SEQ ID NO:6 and/or 7).

Table 3: Human Osteocalcin Regulatory Region

1 ttctccgtc cgatgcgca gggcagggtt gaccgtcgag ctgcacccac agcaggctgc
61 ctttgttgc acaccgggtt aacggggca ttgcgaggca tccccctt gggttggct
121 cctgcccacg ggcctgacag tagaaatcac aggctgttag acagctggag cccagctctg
181 cttgaacctt ttttaggtct ctgatccccg cttccctttt agactcccctt agagctcagc
241 cagtgtccaa cctgaggctg ggggtctctg aggaagagtg agtggagct gagggtctg
301 gggctgtccc ctgagagagg ggccagaggc agtgtcaaga gccgggcagt ctgattgtgg
361 ctcaccctcc atcactccca gggcccccctg gcccagcagc cgcagctccc aaccacatata
421 cctctgggtt ttggcctacg gagctgggc ggtatgacccc caaatagcccc tggcagattc
481 cccctagacc cggccgcacc atggtcaggc atgccccctcc tcatcgctgg gcacagccca
541 gagggtataa acagtgttgg aggctggcg ggcaggccag ctgagtcctg agcagca
(SEQ ID NO:3; , GENBANK™ Accession No. E13404)

Table 4: Mouse bone sialoprotein promoter region and cDNA

1 tctagaaaac actgtccctt taaaatcatt caccacctct ggctccata atctccctgt
15 61 cctcccttcc acacagatcc ctgaggccttg aggagagggc tgtgataaat catccccctt
121 ggagttagca gtcgtaaatgc tctcattctc catgcactgt ctattccgt cccgcgggat
181 tcagtttattc gtgggtgcga gggggaccac gaacctggaa ggaaatgggaa ggaaaagaaaa
241 gagagcggac gaccaagtag attgaacata tcaaggctc gtttattagg ctgaggtgc
301 ttcttttaa agcatacatc acgggaaata tgggaggggt cgagggagaa ttatacaaag
20 361 aacaaagaag tgggcattcg ctgacatggg gccgaaatgc aggcccccagg cagcggcac
421 tctggattt atctctggaa cattgatcct ccttgacagc ctgggggtc aggctggct
481 caggcgtaac tcatgtccctt ggtggcactg ggaactcagg aagagatagg gaagagggga
541 ctataattca gctttacag ctcagggtgc caagaaagga atagggagga aggggggtga
601 taaccagctc ttgtacaag gccatttggc ctgttagggaa gattgtgaag ggctcactt
25 661 ctcacggat ggtctctgac actgtctggc tgggtgtctc cccatctact gcaagactgg
721 gctttctga tgaagtgtaa gccttagtgag ggtgccctgt tcattagaag tcatttgca
781 gtcactcagc agaatattag tagtgggtt ctgtccccct gagagctcac aaccgtctca
841 gtcctgggtt cttagcaccg tgaataattc tatttcaga agttaacatc ctccctca
901 gacaccttg aagcttgtgg gtgttgggt ttctgtgccct tctacctgca cgtctctcca
30 961 tacccaaactg tgagcatttg aaagcgtgtg cttagatgttcc ttgttagct ccccatgtcc
1021 tataaaacac ttgggttgg tagagaactg agcagttcaa acttgcctca actgagctta
1081 tgggggtgaa ttgaatacaa gcaaataaaa ggagcttatt caacttctctt ttgtgggtc
1141 tctattttat tttaaatgc tgaaatactt ttcttagct aaatcatcg aagaatctaa
1201 cagagtcaactt actctggcaa caatactggaa caacaatggc atttattgtat ttctgtaaag
35 1261 tagaagtcaaa cagagaagaa tatggggata aagaatatacg ggataaagaa gacaaccaac
1321 cagagctccc agggcttaaaa ccaccaacca gggagtgacac atggagggac ccatggctcc

1381 atctgtatat gtagcagagg atggcctagt ccatcatcaa tgggatgaga ggcccttggt
 1441 cccatgaagg cctgatatcc cagtgccgg gaatttgggg gcagggagga gagagtggat
 1501 gggtaggtgg gggAACACCC tcatagaAGC aggaggggg gtgggatagg gggTTTGGG
 1561 gtgtggaaat tggaaaggaa gataaacctt gaaacgtaaa taaataaaat atccaataaa
 5 1621 aaaatctctt gggaaagaaaa agatatacaa aatacaaagg cagttccct tgcaactta
 1681 gggaaatgttc agtttgcCAA tgcATGcAgT aagtttattt tccAGTAATTt attcaataac
 1741 catgaACTgc tCTCTGGcAg tgCTAGTAATt tattctcac tcatAGGAaaa aaaattacat
 1801 aagaAGACGA ctAGAAATAA gattatacGA tGTGcAGTGG cCTCATTAC acAGCAAAAGG
 1861 gCCACATAGG ggataatccc aaggacttgt tCTATGAAAG gttACATCAG CTCCTGGTC
 10 1921 tcaACCTCGA acgCTGtaAC gttCACAGTC agcATTGTC ttAGCAAAAG CTTAGGTAAT
 1981 ctgACTGGTT taataatATC agtttGACT tacaAGCCTC tGAATATGT ttcAGGGAGA
 2041 aatataAAAGG aatcaatATT aaACTATCTC ttggCATCAA CTCATTCTC aattcAGTAC
 2101 ttttagACCC atGCAGTGCT GTGTGAAAGC cAGCTTCCT tCTTTCAAC acAGTGAaaa
 2161 CCTGTATCAT tGTGAAAGCT tAAATGCTTA agtCTTTGC tattttttt atttGAAATG
 15 2221 cAGTATATTAA tTATATATAT tCAGAACTCT AACTACCACt TTCTCCTCAC CCTTCATTA
 2281 aatccccacAA tgcaAGCCTC ttggcAGAAG gcccACCTTt catgtttattt caACTGAGGC
 2341 tgaATCTGA aatATGTTG aagtTTGGG tCTCTGGTG agAACCCACA gCCTGACGTT
 2401 gtGCTGGCCA cAGCTGTGAT tggcgttGA gaggcGGAGA aggGTTATA gTCAGCAAGA
 2461 gcaAGTGAAT gAGTGAATGTA cAGCCGGAG aacaATCCGT gCCACTCACT CGACTCGAGC
 20 2521 caaggacTG gCCGAAAGGA aggTTAAGGT aatGGGCAAG gACCTCACAG CCAGGTAATG
 2581 ggcAAGGACC tcACAGCCAG gCACCTAGT CTCCCTGTG tggCTTGGC tggAGTTG
 2641 tagtgcAGC atggatCTTA ctgcACAGTG cacAGTGGCT ctAGTGAAC tttGCTTGC

(SEQ ID NO:4;; GENBANK™ Accession No. AF071079; promoter region,
nucleotides 1-2472)

25

Table 5: Rat bone sialoprotein promoter region

1 aagCTTAGGG aacATTcAGC ctGCCAACAT acGCGGGAAg tttatTTCC agtGATCCT
 61 tcaATGGCCG tggAACTGCT ttCTGGCAGT gCTAGTAATTt CTTCCTCCT cAGAGGGAAA
 121 gataCATAGG aAGAGGACTT agAAATAAGC ctGAGAGTAT acAGCGCTG atGACCTCAC
 30 181 tcGCACAAACG aAAGGCCATG tCCCGATGA tgCCAACTAC ttGTTGAT gagAGTTAA
 241 tcAGCTTCTT ggtCTGAGCC tcaaATGTG tagCTTCAC agTCAGCACA gttAGCAAG
 301 cCTTGGCAGC cCGGCTGGCT ttACAATACT gattCTGACT tacGAGCCTC tGAATGCA
 361 ttcAGAAAGG aAtATAAAGG gatCTTCACT gaACACCTCT tGTCATCAAC tCGTTCC
 421 attCAGTGCT ttAGGCTCG ggcAGTGCTG tGTTAACAG aggCTAGTTt tCCTTCTTT
 35 481 caACATAGTA AAAACCTGTA tcATTGTGAA agTTAAATG ctaAGTGTG tGCCCATT
 541 gttATTGTA aAtGAGTGT ATTATTAAG atATCAGAA ctCTAACTAC catCTTC

601 tcagccctca attaaatccc acaatgcgac ctctggcag caggcgcc ttcatgtt
 661 attcaactga ggctgagtct taaaaacgtg ttgtatcac ggattttctg gtgagaaccc
 721 acagccgtac gtgcacccgg ccgtgaccgt gatggctgc tgagaggaga agaagggtt
 781 ataggtcagc aagagcgagt gaatgggtga gaggcagccg ggagaacaat ccgtccact
 5 841 cactcacttg ctctctccag ccaggactgc cgaaggtaag gtaatggcc agcacctcac
 901 agccacctgc ctaggcttc ctgtgtggct ttggcttggaa atttgtcgtt gaagcatgaa
 961 tcttactgct tgggcaccaa tggctctggt tgaacttttag cttgctgtga aatgggacct
 1021 ctgagtttag ttcttcca aagaccaggc tggtaacgt aagcatgcg taaaactgct
 1081 tcagatttgtt acc

10 (SEQ ID NO:5; GENBANK™ Accession No. LO6562)

Table 6: Mouse dentin sialoprotein gene regulatory regions (5' to exon 1)

1 gaattcttt cccattggta acgtaaaaga ccactactta attgagtttag cttaggctca
 15 61 acaaacagac ttataacaac ttaacttctt tcacattt gaaaaattaa tcagtatcg
 121 cactgagaag gcagaaacag gttagactcc atgagttca gcccggctg atctacatag
 181 gaattcttagg acaaggcaggg cttagtagag ataccctatc taaaaaaaaacc aaaacccaaa
 241 aacattacgt ttaaggcagat ttatgttga ccctaaatgt ttgtcttagt gaaggccc
 301 aatgcttta gcaaatagtttt ctttgttagt ttggagatgt ttgtgtgcta atacagctat
 20 361 caagcacttc tgtagagaca ccgaagatct tcttaactct ccatcaggctc tggagagctg
 421 ttcaaatctg ctattacaac caagtttagga agaggaaggc aattccttag gaaagtggca
 481 ttcttaataa tggatggccc tttaagatgc tcaaagaacc aagaaccatg cagtgtaaat
 541 aatagcaaag tgttactat ggaagtgcag cttcgaggaa actcccttcc tatcactgga
 601 acctgtccaa tccctaccta catgaatatg ttgttaattt ctctcgtat aaagctctga
 25 661 agatgtgtt gctggatagt gatataat ttctgtatcat atgtgtttga catcttcag
 721 tagtgtgaca taaaaacatg gacacatccc taagctggta cacagagact ccaattggct
 781 agtgtggcgc tcataagcta gagaaatggc tcaggatca tcttgtatcc ccaaggctcg
 841 agagaatgtt gggtcaggc aagtacit tcccttctgg aagcacagcc tgtttcctt
 901 ttctgtactc tataatgtt acatataatg gagcaaaagaa tggaaatgtt gtctgtgg
 30 961 tggatgtgtg tgcaactgtt actacgcattt agataccctt caccatgtt cacccttgg
 1021 acagctttt ttaaattttt ttgttattaa attaatatgttataaaagaaaa aacccaaaac
 1081 ctttatgtca gtgttagat taaaatcgaa aggtttccctg aagttactgt ttataaattt
 1141 tttaaagat cccttaggca gtgtcaagac tggatgtcatgc ggacagccgc ttgaattata
 1201 ggcaccaac ttaaatatgtt acctcaggaa tgatagggtt cttaatgtc cagtcgtatt
 35 1261 tactagagaa accttagat ttcttagatt gccgaccaa gcaagaggag aaatgcagg
 1321 tgacagagtc taagtggc tttcagata tttcacactg attatctata ttaagacac
 1381 aaaaacgtct tccaggagctt atttaattaa gtgaaaatgtt gtctgtatcc ttggaaacca
 1441 aagggtctc tagccaaacg taccggcagcg cgagggatgtt gggcgatattt acagccat

1501 aggacacactg actcttaaa ccccccacatc agggatccta agcagtgtt ggttgagaaa
1561 attatcaaac tgaatttaaa ttccaggcagg tacaaaatgg tcacgcaaaa agcccgaggac
1621 agtgtgc (SEQ ID NO:6; nucleotides 1-1627 of GENBANK™ Accession No.
AJ002141).

Table 7: Mouse dentin sialoprotein gene regulatory regions (intron between exon 1 and exon 2)

DRAFT GENBANK

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3541 ctcaactctt ggccctgggg ctttgtctca aggcacccca ttttagttt tagaataatt
3601 gaagggaccc cagctttct tagtttcac ttgacagcta taaggaaggg tgaagcatct
3661 tttcagaga tcctagaatt gtgttctcac ttctgtcaag taataaaacaa tatatattca
3721 ttgtatgtttt attctattcc cctattaacc ttggattttta atcaaggaca ttttatgtg
5      3781 tgcaagggtgg taatcattaa ttcttgga aggtcacaag ataggagaaa acaattctt
3841 ctatagtaaa acaccatgtat acaaataaaat ttagtttttag aaaaatggaa cctgaagttt
3901 tgattcacat agattttat agtttacag gctccattcc aatgtatgaa aaatatgtat
3961 ctgattctgt gaatttgcattt tgcaaagggtt gaaagatttc actctgaag cctctctc
4021 tcagctcctc cctcagtccg agactgcata gtgcccgggt aagggtgggg tgtcccttgc
10     4081 cctcaggagt gcttgtcg cagcaggctc tgcaagggtga ccttgcattt gctcagaaga
4141 cactgatgtat caagatgtcg cgctgggctc cgagacctga tgccagttag gaggaagatg
4201 gggtagctag gcaacttcaa aacagtgcaa tgtgctgcca gcatcgagcg agcggagggt
4261 gcacaaggctg atgcgtgtg aggaaggggag ctaaaagatgc cttcagaaaag cttttgggg
4321 gtgattcttc tgccaaccccc taggatattt tgagctacag agttattaaa ccagactgag
15     4381 gaaacaaaaag cccaataaaag ctattgaaag tgcccaagct cagagagcag atagcagggg
4441 aaggatttga attcagggtt ctgaaaccaa atcctgtt ctcttcctca gcctaaactc
4501 tctttccctt aaacactgtat agaggaagat ttcttcctt tactggata acgcccattt
4561 ctatatacgtat caggtggaa attacaagtgtt tttatcatt tacaatctac ttttagttaa
4621 tgatgctaa agctagccca ggagagacgt taccctcatg gataacagca tagggccaga
20     4681 gccacgagct atgtactctg tatcttcattt gctgttgctt ccacaggcag gttaggtcag
4741 aagccatgac agtccctgagc atgcagaggc ccccacatac ccagggttat ttctggaaacc
4801 tgggtgtttt tctcacattttt gtacttttcctt cttgtccttag aaaaggccca aatgtaaac
4861 caaaaatatttgggtactgtg gctgtcatctt ttcattttt gacccgtttt gtgggtttct
4921 ttgttctaaa cag (SEQ ID NO:7; nucleotides 1-1627 of GENBANK™ Accession
25     No. AJ002141)

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Expression of the nucleic acid is preferably inducible. Osteoblast or odontoblast transcriptional regulatory DNA is used to control expression of IL-4 or another anti-inflammatory polypeptide in a transcription unit. A truncated fragment of such promoters, e.g., containing part of SEQ ID NO:3, 4, 5, 6, or 7, which functions to preferentially direct transcription in odontoprogenitor cells or OPCs (compared to other cell types) may be used. The regulatory sequence, e.g., a cis-acting cell-specific transcriptional regulatory element, is positioned 5' to a heterologous nucleic acid sequence, in a transcription unit. All or part of one of the nucleotide sequences specified in a reference sequence, its complementary strand or a variant thereof may be used in to direct transcription of a heterologous nucleic acid sequence. A nucleic acid fragment is a portion of at least 20 continuous nucleotides identical

to a portion of length equivalent to one of the reference nucleotide sequences or to its complement.

Expression of a heterologous polypeptide-encoding sequence is regulated by contacting the cells of the invention with an antibiotic compound such as tetracycline or a 5 tetracycline analogue (e.g., minocycline or doxycycline). For example, tetracycline is systemically administered at least 2 days before periodontal surgery, e.g., and the time at which cells of the invention are implanted, and/or for at least 2 days after surgery and/or implantation. Expression of the heterologous polypeptide by the cells is turned on while the antibiotic is present in the tissue, i.e., while it is being administered to the cell implant 10 recipient. Expression of the recombinant anti-inflammatory polypeptide decreases and ceases after administration of the antibiotic is stopped. Typically, an antibiotic administered 8-12 days prior to surgery and 8-12 days post-surgery. Similarly, antibiotics are administered before and after orthopedic surgery, e.g., surgery for cartilage removal from articulating joints or for removal of metastatic bone tumors (at which time the cells are 15 implanted at or adjacent site to diseased tissue). The cells may be implanted before, during, or after implantation of a dental orthopedic prosthesis. To treat advanced periodontal disease, the cells are administered locally to the periodontal ligament in the mandibular section of the jaw. A clinical benefit is conferred by using the cells to inhibit osteolysis in a mammal, e.g. a human patient, that is suffering from or at risk of developing periodontitis or 20 other bone disorders which may lead to bone loss, e.g., alveolar bone loss.

The methods described herein are also applicable to veterinary use, e.g., to treat dogs, cats, horses.

The invention includes OPCs which are genetically modified to contain a nucleic acid encoding an anti-inflammatory polypeptide. OPCs are derived from bone marrow stromal 25 cells and have been differentiated *ex vivo* in the presence of ECM. As is described above, the OPCs preferably contain a nucleic acid encoding a cytokine such as IL-4, or an agonist thereof, operably linked to a promoter which directs transcription of a nucleic acid to which it is linked preferentially in cells which have differentiated into osteoblasts.

For treatment of bone disorders, the cells are implanted into the bone marrow of a 30 recipient mammal or into an articulating joint of the mammal. For example, the cells are administered intratibially or intrafemorally. The cells are implanted locally, e.g., at the site

of bone loss or adjacent to such as site, e.g., in the bone marrow, and expression of the recombinant polypeptide by the cells is regulated by systemically administering an antibiotic such as minocycline or doxycycline. Methods of transplanting cells into the bone marrow of a mammal are well known in the art, e.g., as described in U.S. Patent No. 4,188,486. The dose
5 of cells to be administered ranges from 1×10^5 cells to 1×10^{10} cells in volume suitable for the location of transplantation (e.g., a smaller volume is used for implantation into mandibular tissue or into the periodontal ligament compared to implantation into the bone marrow of the femur). Clinical protocols for such implantation procedures are known in the art. For example, a dose of 1×10^8 cells per kg of body weight is administered to femoral
10 bone marrow. Repeated implants may be required in the case of long term diseases such as rheumatoid arthritis.

Inhibitors of cyclooxygenase II (COX-2) or tumor necrosis factor-alpha (TNF α) are optionally administered. COX inhibitors include aspirin, ibuprofen and indomethacin, as well as bisaryl COX-2 inhibitory compounds (e.g., as described in U.S. Patent No.
15 5,994,379) and (methylsulfonyl)phenyl-2-(5H)-furanones (e.g., as described in U.S. Patent No. 6,020,343).

The isolated genetically-modified OPCs are used to treat individuals suffering from or at risk of developing a bone loss disorder such as rheumatoid arthritis, osteoporosis, periapical or endochondral bone loss, artificial joint particle-induced osteolysis, bone fracture
20 or deficiency, primary or secondary hyperparathyroidism, metastatic bone disease, osteolytic bone disease, post-plastic surgery, post-prosthetic joint surgery.

By the term "osteoprogenitor" is meant a differentiated bone precursor cell derived from a bone stromal cell. By the term "odontoprogenitor" is meant a differentiated bone precursor cell derived from periodontal ligament. The differentiated state of the bone
25 marrow stromal cells or ligament derived cells is induced by culture in the presence of ECM. Preferably, the cells are cultured in the presence of a BMP such as BMP-2, 4, or 6. Differentiated progenitor cells have enhanced ability to build bone tissue, compared to undifferentiated stromal cells. OPCs or odontoprogenitor cells are distinguished from bone stromal cells (as well as fat, muscle, or cartilage cells or tissue) by the production of alkaline
30 phosphatase, expression of osteocalcin, and expression of bone sialoprotein (in addition to the expression of dentin sialoprotein in the case of odontoprogenitors).

The *ex vivo* cell-based therapeutic methods of the invention has several advantages over standard gene therapy protocols. For example, the cells expressing the recombinant anti-inflammatory polypeptide are isolated, i.e., purified from cells which do not have the desired phenotype. A population of isolated OPCs or odontoprogenitor cells is at least 75%,
5 more preferably 85%, more preferably 90%, more preferably 95%, more preferably 98%, more preferably 99% or 100% OPCs or odontoprogenitor cells, respectively.

DNA is introduced into isolated cells *ex vivo*, thus avoiding or minimizing the possibility DNA uptake by non-target cells in the body. Another measure of safety is conferred by using a transcriptional regulatory element and a promoter that directs
10 transcription only in the isolated cell type. *In vivo* expression of the recombinant polypeptide is further regulated by the systemic administration of an antibiotic or antibiotic analogue.

OPCs are isolated and expanded from stromal cells from bone marrow aspirates, and autologous bone marrow stromal cells are expanded. The cells are optionally frozen and stored in liquid nitrogen for long periods of time before being differentiated and transduced.
15 These "banked" autologous cells allow for multiple inoculations over a long period of time, which is advantageous since RA may persist for many years

Other features and advantages of the invention will be apparent from the following detailed description and from the claims.

BRIEF DESCRIPTION OF THE DRAWINGS

20 Fig. 1 is a bar graph showing alkaline phosphatase acitivity of rabbit bone marrow stromal cells plated on ECM produced by untransduced C3H10t1/2 cells, or C3H10T1/2 cells transduced with and without BMP-6, or on plastic alone. Activity is expressed as $\mu\text{mol p-nitrophenol produced/min/mg protein} \times 10^4$ at days 1 and 21.

Fig. 2A is a diagram of a periodontal ligament biopsy.

25 Fig. 2B is a diagram of the method for differentiating odontoblast precursor cells on an osteoinductive matrix and transduction of the cells with a regulatable therapeutic gene.

Fig. 2C is a diagram of a device for implanting cells adjacent to teeth in need of therapeutic intervention.

30 Fig. 3A is a photomicrograph of rabbit bone marrow stromal cells cultured for 21 days after plating on tissue culture plastic. Magnification, 100 \times .

Fig. 3B is a photomicrograph of rabbit bone marrow stromal cells cultured for 21 days after plating on ECM-coated dishes from a C3H10T1/2 cell line transduced with rhuBMP-6. Magnification, 100x.

Fig. 4 is a bar graph showing the effect of rhuIL-4 on rhuIL-1 alpha-induced PGE2 by 5 rabbit osteoprogenitor cells.

DETAILED DESCRIPTION

Cell-based delivery of rhIL-4 at the site of an osteolytic lesion allows rhIL-4 to be concentrated near an inflammatory site where inflammatory effector cells, e.g., macrophages, and 10 osteolytic effector cells, e.g., osteoclast precursor cells, are located. Adverse effects of rhIL-4 on thymocyte and T cell function are greatly decreased since the cytokine acts locally as opposed to functioning throughout the body when delivered systemically.

The cells described herein are committed to the osteoblastic lineage. Differentiation was induced by stimulating the bone marrow stromal cells to differentiate by exposing the cells to an extracellular matrix such as Matrigel (Becton Dickenson) or other commercially available matrix 15 preparation in the presence of bone morphogenetic proteins. This step is typically carried out before the differentiated cells are transduced with retroviral expression vectors containing genes encoding one or more therapeutic proteins. This is advantageous in that the transduced cells cannot differentiate into cartilage, muscle or fat cells once implanted *in vivo*. In contrast, the pre-osteoblastic pluripotent bone marrow stromal cells still retain the potential to differentiate.

Unlike the retroviral vectors employed in the prior art which use viral promoters, the 20 retroviral expression vectors in the OPCs have been constructed to use osteoblast specific promoters to initiate transcription of the reverse transactivator Tetracycline Activator (rtTA) gene, which in turn regulates production of the rhIL-4 therapeutic protein. This approach provides the advantage of increased safety because the osteoblast promoters direct transcription more efficiently 25 in OPCs (compared to other cell types such as stromal cells) and are far less likely to be inactivated *in vivo* than viral promoters. For example, the viral vectors contain a doxycycline-inducible system which regulates the expression of the interleukin-4 encoding sequences.

The OPCs described herein were modified to increase the expression of the alpha-5 30 integrin receptor. This modification allows the cells to adhere to bone matrix proteins when implanted *in vivo*, which confers the added advantage that the OPCs may be inserted directly

into osteolytic sites without prior encapsulation, e.g., in porous calcium phosphate ceramic cubes or other types of encapsulated devices.

The cells are used for autologous or allogeneic cell transplants to serve as a cell-based platform to deliver the recombinant human interleukin-4 (rhIL-4) protein in a site-specific, 5 regulated manner. RhIL-4 acts on defined cell types when the OPCs are implanted at the site of osteolytic bone lesions. The OPCs, genetically-engineered to produce rhIL-4 are used to i) promote tumor reduction when used in conjunction with anti-cancer drugs, ii) inhibit formation of osteoclasts which resorb bone, and iii) stimulate new bone growth. The methods result in improved clinical outcomes. The OPCs, engineered to secrete rhIL-4 may 10 are implanted in patients who are undergoing revision of a artificial joint replacement due to the development of implant-induced osteolysis, as well as in patients suffering bone loss due to rheumatoid arthritis and in the oral cavity due to severe periodontal disease.

IL-4 and agonists thereof

IL-4, an anti-inflammatory cytokine produced primarily by Th2 cells and 15 macrophages exhibits anti-inflammatory and immunosuppressive properties. Bone-derived cells, e.g., differentiated osteoprogenitor cells or odontoprogenitor cells, which are genetically-altered to produce recombinant human interleukin-4 (rhuIL-4) are administered to diseased bone tissue. The invention provides a gene therapy approach to deliver rhuIL-4 locally at inflamed joints by targeting bone marrow stromal cells that have undergone 20 partial differentiation with a viral plasmid expression system containing a cell-specific promoter. In order to prevent potentially harmful effects of rhuIL-4, due to high local or systemic concentrations, the production of rhuIL-4 is regulated by the oral administration of antibiotic analogues. Articular cartilage degradation and bone resorption, associated with rheumatoid arthritis, is reduced significantly by the local, regulated release of IL-4 near the 25 site of tissue damage. The mechanism for reduced cartilage degradation and bone loss is based on the ability of IL-4 to inhibit TNF-alpha, IL-1, and PGE2 production, as well as the ability of IL-4 to decrease angiogenesis. Regulated, local release of IL-4 decreases cartilage and bone destruction *in vivo*. IL-4 also has immunosuppressive properties; in situations in which the implanted cells are allogeneic (rather than autologous), the IL-4 produced by the 30 implanted cells may obviate the need to administer systemic immunosuppressive drugs to combat tissue rejection.

Example 1: Differentiation of rabbit bone marrow stromal cells on a BMP-6 containing osteoinductive extracellular matrix derived from C3H10T1/2 cells

Over 185,000 spinal arthrodeses are performed in the US each year, with non-union rates as high as 35% reported in the most commonly performed procedure, posterolateral lumbar intertransverse process fusions. Autologous iliac crest bone is the gold standard graft material, but it is limited in quantity, and the morbidity of harvest is not insignificant. The data described herein indicate that BMPs induce differentiation of multipotential stromal cells from rabbit bone marrow an osteoblastic lineage. The stromal cells are exposed to ECM secreted by a transfected murine cell line constitutively expressing or overexpressing mRNA for BMP-6.

Cells from the C3H10T1/2 murine fibroblast line were transduced with either an LXSN vector containing the rhBMP-6 gene or the same vector without the gene. The cells were cultured under standard conditions in DMEM (Gibco; Gaithersburg, MD) supplemented with 100 units/ml PCN, 100 µg/ml streptomycin, and 10% FBS (Hyclone; Logan, UT). Four days after reaching confluence, the cells were lysed sequentially with water and 0.1% Triton X-100. The plates were gently washed with phosphate buffered saline (PBS), leaving the extracellular matrix and the BMP-6 protein adherent to the plastic dish.

Bone marrow was aspirated from the femurs of two New Zealand white rabbits and suspended in DMEM with 100u heparin/cc. The cell suspension was diluted with PBS, 2%BSA, 0.6% sodium citrate, and 1% penicillin/streptomycin. The suspension was then layered on a Ficoll-Paque (Amersham Pharmacia Biotech; Piscataway, NJ) gradient, and centrifuged at 600 × g for 20 minutes. The cells at the interface were isolated, washed, and re-centrifuged at 500 × g twice. They were then cultured to confluence in a T-75 flask under standard conditions in a-MEM with L-glutamine 2mM, without nucleosides, supplemented by 12.5% FBS, 0.2mM I-inositol, 20nM folic acid, 0.1mM beta-mercaptoethanol, and 1% penicillin/streptomycin. The cells were re-plated in triplicate on the ECM produced by untransduced cells, LXSN transduced cells, or LXSN-BMP6 transduced cells, or on plastic, and cultured for 21 days.

Alkaline Phosphatase (ALP) activity was determined on day 1 and 21 after plating of stromal cells on ECM. The plates were scraped and rinsed with 0.5M CAPS, pH 10.5, and

sonicated. 0.5ml of 0.5% p-nitrophenyl phosphate was added to each sonicate, and incubated at 37° C for 30 minutes. 0.2M NaOH was added to stop the reaction, and the amount of p-nitrophenol produced was determined by spectrophotometry at 405nm. ALP activity was expressed as μ mol p-nitrophenol/min/mg protein. Protein content was determined by
5 Bradford protein assay (Bio-Rad, Hercules, CA). As an additional control, the ECM produced by each C3H10T1/3 cell line was also assayed for ALP activity at day one, and after exposure to media for 21 days.

The ALP activity in the ECM alone was negligible at day 1 and 21. The activity from each of the ECM-exposed marrow cells was likewise negligible at day 1, as was the activity
10 from the cells on plastic. However, at day 21, there were striking differences (Fig. 1). The ALP level produced by marrow cells plated on plastic was unchanged. That of marrow cells plated on the ECM from untransduced or LXSN cells each increased 400%, while that of the marrow cells plated on the ECM from BMP-6 transfected cells increased 700%.

ECM-bound BMP 2 and 4 produced by neonatal mouse calvarial cells stimulated
15 ALP activity in mouse bone marrow cells. Exposure of stromal cells to EMC in the absence of BMP-6 increased ALP production, presumably due to the presence of type collagen in the matrix. The further increase of ALP production by BMP exposed cells is due to increased osteoblastic differentiation of the stromal cells.

These results indicate that exposure to ECM-bound BMPs induces stromal cells to
20 differentiate along an osteoblastic lineage. Cells of the osteoblastic lineage, e.g., OPC's or odontoprogenitor cells, are identified and purified by virtue of their expression of marker genes such as alkaline phosphatase, osteocalcin, and bone sialoprotein (in addition to dentin sialoprotein in the case of odontoprogenitors). Probes to detect the marker genes are known in the art (e.g., as described by Guo et al., 2000, Calcified Tissue International 66:212-216).
25 Marker gene expression is detected by measuring transcription of the genes (e.g., using labeled nucleic acid probes in *in situ* assays) or by immunohistochemistry to detect antibody binding to the gene products. The assays described above are used to distinguish stromal cells from OPCs and odontoprogenitor cells.

Example 2: Inhibition of Alveolar Bone Loss by Cell-Delivered IL-4

The osteoclast is responsible for mediating excessive bone resorption during progressive periodontitis. IL-4 inhibits osteoclast differentiation and function. Autologous cells are engineered to express IL-4 and permanently implanted at sites of inflammation, e.g., 5 in the mandible, in soft tissue adjacent to affected teeth, or in the periodontal ligament, using methods known in the art.

Periodontal disease is induced in C3H mice by repeated injections of LPS derived from the clinically-relevant microorganism *Porphyromonas gingivalis*, an art recognized model of periodontal disease. Mice with periodontal disease are treated using C3H10T1/2 10 cells genetically engineered to produce IL-4 in a regulatable manner. Production of interleukin-4 is regulated by providing antibiotic orally, e.g., in the drinking water. Cells are implanted locally, at sites of bone resorption, thereby bypassing the need for either systemic administration or repetitive local injections of a bioactive molecule. Optionally, antibiotics are placed in the periodontal pocket following implantation of cells for periodontal disease. 15 This cell-based approach for local delivery of interleukin-4 utilizes tissue engineering to inhibit resorption of alveolar bone.

The murine molony retroviral vectors used herein are well characterized and are non-immunogenic in humans or mice.

Standard *in situ* hybridization (ISH) is used to detect IL-4 production as well as 20 characterize the osteoclast phenotype in cells that have populated mandibler bone or other bone tissue of cell implant recipients.

Example 3: Gene therapy vector

A tetracycline analog-regulated expression system is used to direct production of recombinant anti-inflammatory compositions. A dual “tet-on” retroviral system is used for 25 the following reasons; i) the vectors are commercially available and the packaging cells produce high retroviral titers, ii) the murine molony retroviral vectors have been well characterized, are non-immunogenic, and have been used in safely in humans, and iii) the use of two retroviral vectors in the “tet-on” mode prevents “leaking”, i.e., recombinant polypeptide expression is extremely low or absent without antibiotic present.

“Tet-off” and tet-on” systems use the antibiotic tetracycline or various analogues to 30 regulate expression. Toxicity of the VP16 viral transactivator fusion protein (tTA) was not

observed and no antibodies were made to the "reverse" (rtTA, tet-on) or rTA, "tet-off" transactivator. The tetracycline analogue doxycycline is the preferred antibiotic inducer for the "tet-on" system and is administered by orally or by intraperitoneal administration using known methods, e.g., as described for mifepristone and rapamycin, as well as by
5 implantation of subcutaneous pellets. Doxycycline and/or minocycline is given orally. Minocycline inhibits the action of matrix metalloproteinases (MMPs) which are involved in breakdown of bone and cartilage. Minocycline-activated site-specific IL-4 production at the inflamed joint acts in a synergistic manner to inhibit inflammation and angiogenesis.
These two drugs may work together and lead to increased benefit for patients suffering from
10 RA. The ability to shut down local IL-4 production by removal of minocycline in the "tet on" gene system is advantageous to prevent deleterious effects of sustained IL-4 production at a site of inflammation such as in a rheumatic joint.

Some viral promoters/ enhancers used in adenoviral and retroviral plasmid vectors are inactivated by interferon-gamma and tumor necrosis factor-alpha *in vivo*. These include the
15 rous sarcoma virus (RSV), simian virus 40 (SV-40), and cytomegalovirus (CMV) promoters. Since levels of IFN and TNF are elevated in RA and OA patients, the use of these standard viral vectors could limit recombinant polypeptide expression, especially if sustained production is required. Given these limitations of viral promoters for long term *in vivo* use, the invention utilizes a constitutive cellular promoter in place of the CMV promoter
20 to control expression of the rtTA transactivator in one of the two retroviral plasmid vectors. A human osteocalcin promoter sequence (e.g., SEQ ID NO:3) is employed to modify a "tet on" retroviral vector for transduction of OPCs *ex vivo*.

Late-stage rabbit osteoprogenitor cells obtained from bone marrow stromal cells were isolated and characterized using marker gene detection. The rabbit OPCs have undergone
25 partial differentiation on a osteoinductive matrix derived from C3H10T1/2 cells that have been transduced with a retroviral vector expressing recombinant human bone morphogenetic protein-6 (rhuBMP-6).

Bone marrow stromal cells are obtained from an individual such as a human patient
8-12 weeks prior to therapy. The cells are expanded, differentiated and transduced with
30 recombinant DNA encoding anti-inflammatory polypeptides *ex vivo*. The OPCs are implanted in the marrow bones adjacent to the diseased or injured site, e.g., in the mandible

or periodontal ligament for periodontal disease or in the marrow of the distal femur and proximal tibia, i.e., in juxtaposition to an inflamed knee joint. This approach allows the rhuIL-4-transduced OPCs to be in close proximity to the bone-resorbing osteoclasts and the pannus/bone interface of the joint. Correct positioning of the implanted OPCs in the marrow
5 is important for the following reasons; i) inhibition of bone resorption by IL-4 is optimized by close proximity of the OPCs to the synovial fibroblasts of the invading pannus and the surrounding osteoblasts, ii) the locally produced IL-4 inhibits osteoclast formation from the differentiated synovial marrow-derived macrophages in the presence of rheumatoid synovial fibroblasts, and iii) the OPC-produced IL-4 locally inhibits neovascularization of the
10 inflamed joint.

Other advantages of using bone marrow stromal cells, which have been partially-differentiated toward the osteoblastic lineage, include a high level of expression of the rhuIL-4 for a given cell population due to the higher number of osteoblastic cells locally present. Moreover, a pure population of differentiated stromal cells is not required due to the
15 high specificity of the osteocalcin promoter for osteoblasts. Adipocyte, muscle, and chondrocytic precursor cells will not express the rhuIL-4 even if these cell types take up the recombinant anti-inflammatory polypeptide-encoding DNA. These features provide a safer and more controlled environment for cytokine release when the transduced osteoblast cells are inoculated into the bone marrow.

20 Example 4: Construction of the rhIL-4 "tet-on" dual retroviral expression vectors

The pRevTet-On and pRev-TRE retroviral expression vectors are publically available from Clontech Laboratories, Inc. (Palo Alto, CA). Both vectors were derived from pLNCX, a retroviral vector capable of producing high-titer virus in the RetropakTM (Clontech, Inc.) PT67 packaging cell line. The core murine Moloney leukemia viral vector for each plasmid
25 consists of a 5' long terminal repeat (LTR) containing a promotor (L) which drives the extended retroviral packaging signal. The remaining DNA consists of pBR322-based plasmid sequences which allow for replication in bacteria and an ampicillin resistance gene for bacterial selection. All plasmids and retroviral expression vectors were purified by cesium chloride/ethidium bromide ultracentrifugation gradients, checked for purity on
30 agarose gels, and analyzed for orientation by restriction enzyme mapping and DNA sequence analysis.

pRev-Tet-On-huOC plasmid construct

The pRevTet-On vector also contained a neomycin phosphotransferase gene, then an internal minimal immediate early cytomegaloviral (CMV) promoter which drives the reverse tetracycline regulatory element (rtTA). A 1.339 kb BamHI/EcoRI cDNA was excised from 5 the pGoscas vector which contains a cDNA of the human osteocalcin promoter. The regulatory DNA was ligated into pRevTet-On digested with BamHI/Clal, which removes the CMV promoter and the rtTA portion of the vector. A 1.05 kb cDNA to rtTA was subsequently ligated into the Clal site of pRev-Tet-On followed by blunt end ligation. This resulted in substitution of the viral CMV promoter/enhancer for the osteoblast specific 10 human osteocalcin promoter in the pRev-Tet-On retroviral vector.

pRev-TRE-rhuIL-4 plasmid construct

The pRev-TRE vector (Clontech), contains a 5' long terminal repeat (LTR) containing a promotor which drives the extended retroviral packaging signal. The transactivator response element (TRE) contains seven direct repeats of the tetO operator 15 sequence upstream of a minimal CMV promoter, which can be bound by the tTA and rtTA transactivators. The rhuIL-4-pCD plasmid was obtained from American Type Culture Collection (ATCC (#57593). A 0.86 kb BamHI hIL-4 insert was isolated and used for subsequent subcloning into the BamHI site of the multiple cloning site in the pRev-TRE vector. Restriction enzyme mapping was performed to check for the correct orientation of 20 the rhuIL-4 cDNA insert. The pRev-TRE-rhuIL-4 vector was subsequently transfected into the PT67 packaging cell line (Clontech), selected with Hygromycin B, and high titer clones were assayed using serial dilutions of viral supernatants before infection and Hygro B selection of NIH3T3 cells. PT67 packaging cells containing the pRev-TRE-rhuIL-4 vector 25 were used to sequentially infect the rabbit osteoprogenitor cells along with the pRevTet-On-huOC vector. The transduced cells are subjected to selection with G418 and Hygromycin B and clones producing rhuIL-4 in response to tetracycline ana-log treatment are isolated. The optimal doxycycline and minocycline concentrations for induction of rhuIL-4 production in the transduced cells is performed using standard methods.

Example 5: Isolation and Osteogenic Differentiation of Rabbit Bone Marrow Stromal Cells

30 Bone marrow aspirate (approximately 1-2 c.c.) was obtained from the greater trocular region of the femur of anesthetized New Zealand white rabbits and suspended in DMEM

with 100u heparin/cc. The cell suspension was diluted with PBS, 2%BSA, 0.6% sodium citrate, and 1% penicillin/ strep-tomycin. The suspension was then layered on a Ficoll-Paque (Amersham Pharmacia Biotech; Piscataway, NJ) gradient, and centrifuged at 600 × g for 20 minutes. The cells at the interface were isolated, washed, and recentrifuged at 500 × g twice.

5 They were then cultured for 2-3 weeks until confluent in T-75 flasks at 37 degrees C in 5% CO₂ in alpha-MEM with L-glutamine 2mM, without nucleosides or glucocorticoids, supplemented by 12.5% horse serum (Sigma; St. Louis, MO), 12.5% FBS, 0.2mM I-inositol, 20nM folic acid, 0.1mM B-mercaptoethanol, and 1% penicillin/streptomycin. The cells were then harvested in DPBS/EDTA /pancreatin and stored in liquid nitrogen in freezing media.

10 Aliquots of BMSCs were subsequently thawed and replated in triplicate on plastic dishes or dishes coated with the ECM produced by untransduced cells, LXSN transduced cells, or rhuBMP6 transduced cells, and cultured for 1 and 21 days. Examination of the thawed cells by phase contrast microscopy revealed that BMSCs cultured on tissue culture plastic for 21 days retained their fibroblastic, spindle-shaped morphology, as shown in Fig. 3A. In

15 contrast, BMSCs plated on the BMP-6-containing ECM for 21 days became more cobblestone-shaped in appearance (Fig. 3B) and resembled osteoblast cells.

Alkaline phosphatase (ALP) activity was determined on day 1 and 21 after plating of stromal cells on ECM. Representative results are shown in Fig. 1. These data indicate that stromal cells may be isolated, expanded, and stored frozen before undergoing further differentiation *ex vivo*. The data also indicate that BMP-responsiveness with respect to alkaline phosphatase induction demonstrates that the BMSCs have undergone partial differentiation to become late stage osteoprogenitor cells

Example 6: Effects of rhuIL-4 on Rabbit late-stage Osteoprogenitor Cells

The effect of rhuIL-4 on PGE2 production by late-stage rabbit OPCs was analyzed.

25 OPCs were harvested from the BMP-6/ECM coated dishes by trypsinization, counted, then replated in 6-well dishes in alpha-MEM plus 1% FBS and 10-5 M arachidonic acid for 24 hr. The cells were then incubated for an additional 24 hr with rhuIL-1 (2 ng/ml) in the absence or presence of IL-4 (25, 50, and 100 ng/ml). PGE2 levels were measured in the conditioned cell culture media by enzyme immunoassay (BioTrak RPN 222, Amersham Pharmacia Biotech, Inc., Piscataway, NJ). As shown in Fig. 4, there was dose-related effect of IL-4 on the inhibition of IL-1-stimulated PGE2 release by the rabbit OPCs.

These results indicate that rhuIL-4 blocks the effect of IL-1-alpha on the induction of PGE2 production in osteoprogenitor cells, an important intermediate step in osteoclast-mediated bone resorption and that IL-4 is beneficial for reducing inflammation associated with RA (since PGE2 is a potent mediator of the pain and edema associated with rheumatoid synovitis).

5 Other embodiments are within the following claims.

What is claimed is:

- 1 1. An isolated odontoprogenitor cell comprising a nucleic acid encoding an anti-
2 inflammatory polypeptide
- 1 2. The cell of claim 1, wherein said cell is derived from a periodontal ligament..
- 1 3. The cell of claim 1, wherein said polypeptide is a cytokine.
- 1 4. The cell of claim 3, wherein said cytokine is interleukin-4 (IL-4).
- 1 5. The cell of claim 1, wherein said nucleic acid is operably linked to an
2 osteoblast-specific promoter.
- 1 6. The cell of claim 5, wherein said osteoblast-specific promoter is an
2 osteocalcin promoter.
- 1 7. The cell of claim 5, wherein said osteoblast-specific promoter is a bone
2 sialoprotein promoter.
- 1 8. The cell of claim 1, wherein expression of said nucleic acid is inducible.
- 1 9. The cell of claim 1, wherein expression of said nucleic acid is regulated by an
2 antibiotic compound.
- 1 10. The cell of claim 9, wherein said antibiotic compound is tetracycline or a
2 tetracycline analogue.
- 1 11. The cell of claim 10, wherein said tetracycline analogue is minocycline or
2 doxycycline.
- 1 12. A method of inhibiting osteolysis in a mammal, comprising introducing into
2 said mammal an isolated odontoprogenitor cell comprising a nucleic acid encoding an
3 anti-inflammatory polypeptide.

13. The method of claim 12, wherein said mammal is suffering from or at risk of developing periodontitis.

14. The method of claim 12, wherein said mammal is suffering from or at risk of developing alveolar bone loss due to periodontal disease.

15. The method of claim 12, wherein said cell is administered to the periodontal ligament in the mandibular section of the jaw.

16. An isolated osteoprogenitor cell comprising a nucleic acid encoding an anti-inflammatory polypeptide.

17. The cell of claim 16, wherein said polypeptide is a cytokine.

18. The cell of claim 17, wherein said cytokine is interleukin-4 (IL-4).

19. The cell of claim 16, wherein said nucleic acid is operably linked to an osteoblast-specific promoter.

20. The cell of claim 19, wherein said osteoblast-specific promoter is an osteocalcin promoter.

21. The cell of claim 19, wherein said osteoblast-specific promoter is an bone sialoprotein promoter.

22. The cell of claim 16, wherein expression of said nucleic acid is inducible.

23. The cell of claim 16, wherein expression of said nucleic acid is regulated by an antibiotic compound.

24. The cell of claim 23, wherein said antibiotic compound is tetracycline or a tetracycline analogue.

1 25. The cell of claim 24, wherein said tetracycline analogue is minocycline or
2 doxycycline.

1 26. A method of inhibiting osteolysis in a mammal, comprising introducing into
2 said mammal an isolated osteoprogenitor cell comprising a nucleic acid encoding an
3 anti-inflammatory polypeptide.

1 27. The method of claim 26, wherein said cell is implanted into an articulating
2 joint of said mammal.

1 28. The method of claim 26, wherein said cell is administered intratibially.

1 29. The method of claim 26, wherein said cell is administered intrafemorally.

1 30. The method of claim 26, wherein expression of said polypeptide is regulated
2 by an antibiotic compound.

1 31. The method of claim 26, wherein said antibiotic compound is tetracycline or a
2 tetracycline analogue.

1 32. The method of claim 31, further comprising administering minocycline to said
2 mammal.

1 33. The method of claim 30, wherein said antibiotic compound is administered
2 systemically.

1 34. The method of claim 26, further comprising administering an inhibitor of
2 cyclooxygenase II (COX-2).

1 35. The method of claim 26, further comprising administering an inhibitor of
2 tumor necrosis factor-alpha (TNF α).

1 36. The method of claim 26, wherein said mammal is suffering from or at risk of
2 developing rheumatoid arthritis.

1 37. The method of claim 26, wherein said mammal is suffering from or at risk of
2 developing periapical or endochondral bone loss, artificial joint particle-induced
3 osteolysis, or osteolytic bone metastases.

1 38. A method of inducing differentiation of a bone marrow stromal cell,
2 comprising contacting said cell with bone morphogenic protein-6.

ABSTRACT

The invention provides compositions and methods to deliver an anti-inflammatory composition, e.g., recombinant human interleukin-4 (rhIL-4), to build (or rebuild) bone tissue. The composition is produced from living osteoprogenitor cells (OPCs) or odontoprogenitor cells.

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FIG. 1

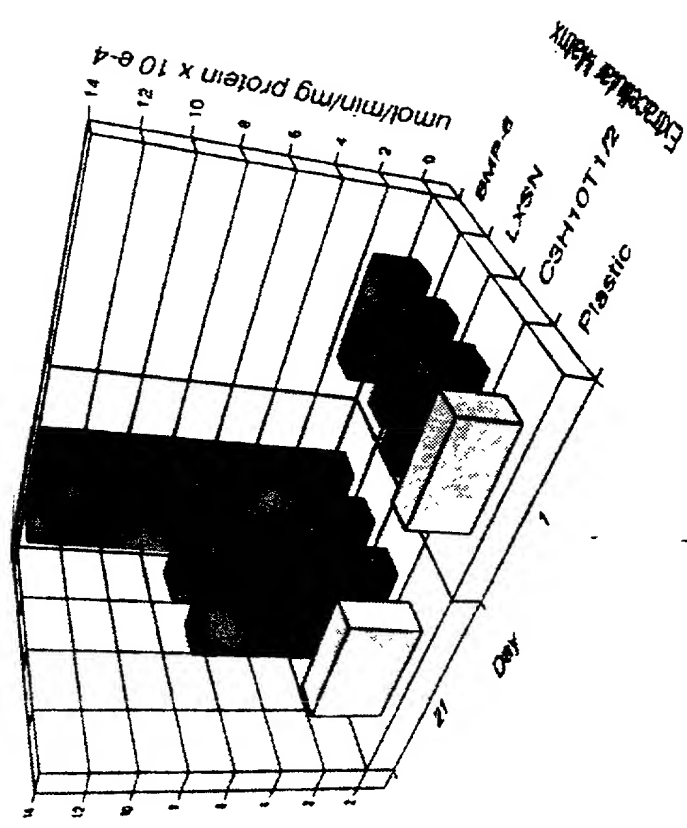


FIG. 2A

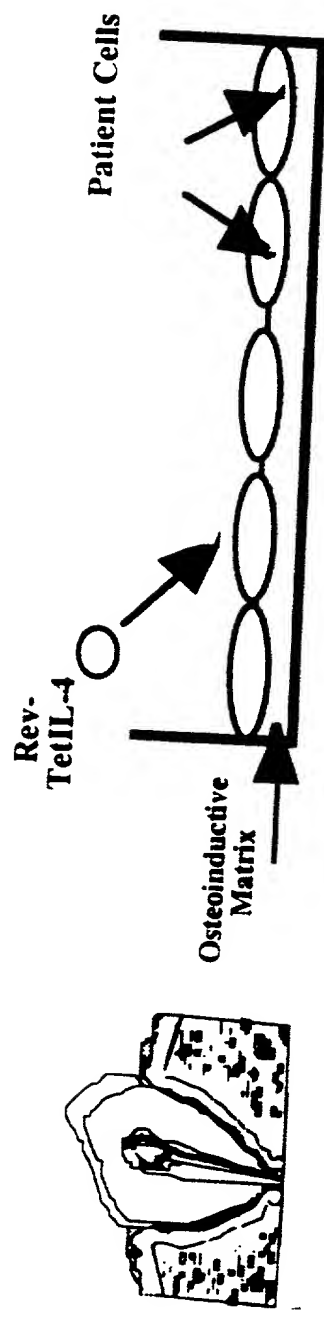


FIG. 2B

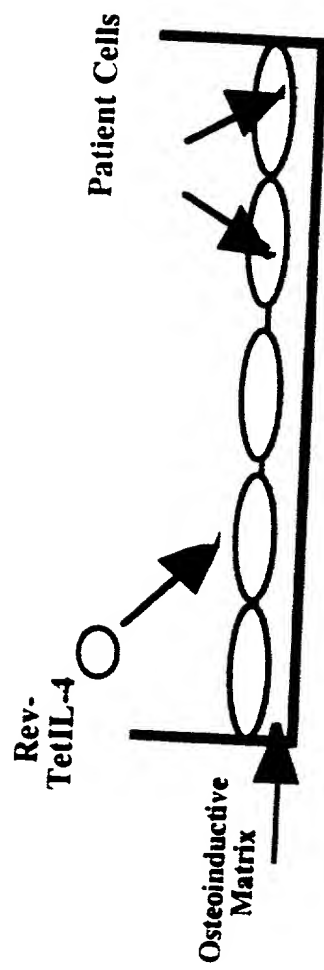


FIG. 2C

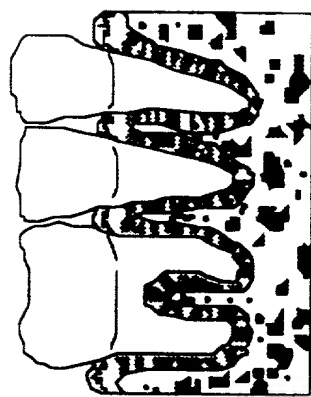


FIG. 3A



FIG. 3B

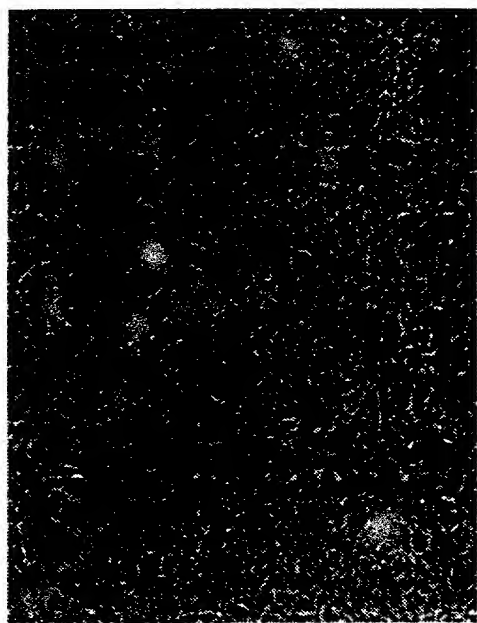
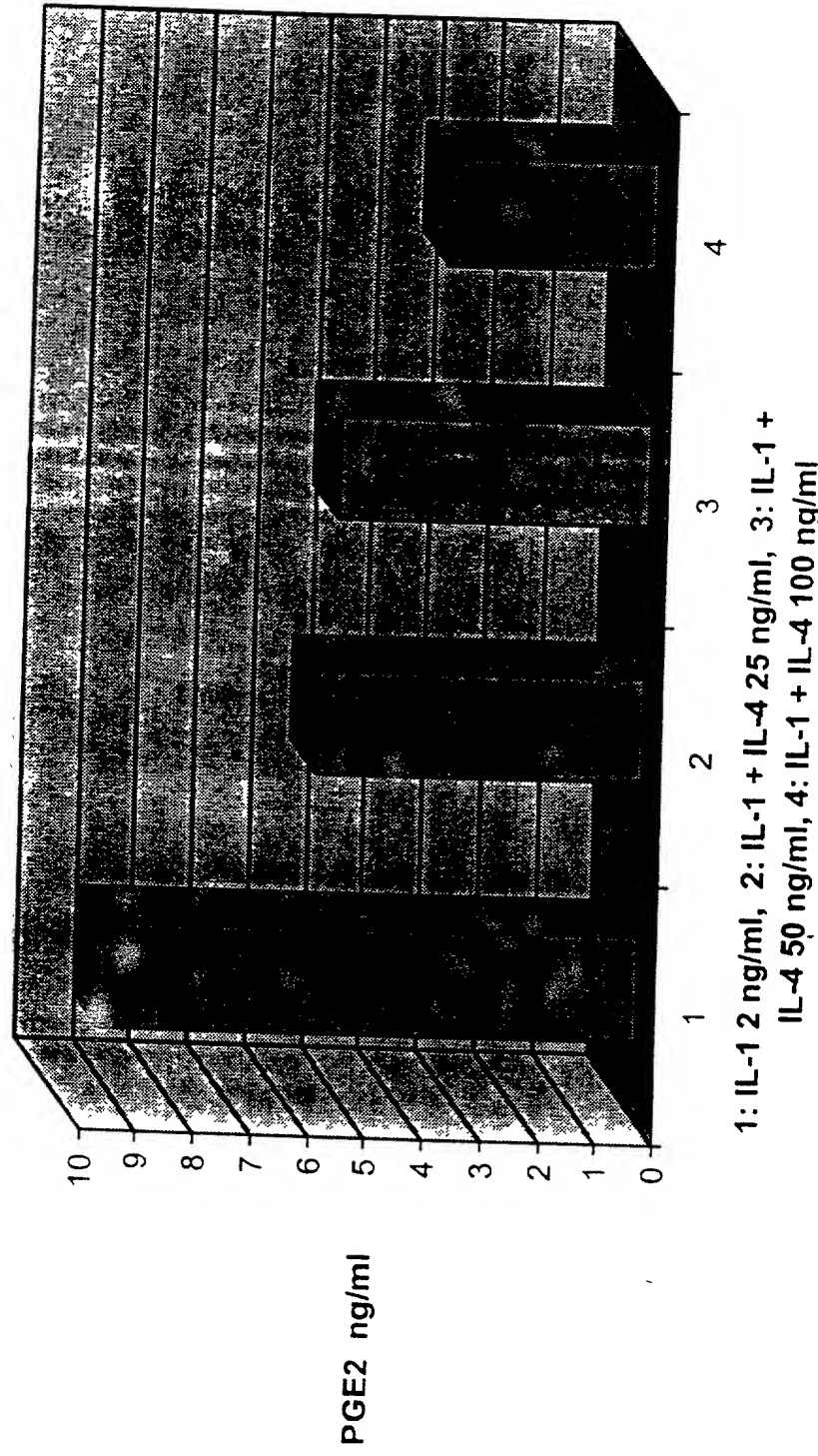


FIG. 4



COMBINED DECLARATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled TREATMENT FOR BONE DISORDERS, the specification of which:

is attached hereto.

was filed on _____ as Application Serial No. _____ and was amended on _____.

was described and claimed in PCT International Application No. _____ filed on _____ and as amended under PCT Article 19 on _____.

I hereby state that I have reviewed and understand the contents of the above-identified specification, including the claims, as amended by any amendment referred to above.

I acknowledge the duty to disclose all information I know to be material to patentability in accordance with Title 37, Code of Federal Regulations, §1.56.

I hereby claim the benefit under Title 35, United States Code, §120 of any United States application(s) listed below and, insofar as the subject matter of each of the claims of this application is not disclosed in the prior United States application in the manner provided by the first paragraph of Title 35, United States Code, §112, I acknowledge the duty to disclose all information I know to be material to patentability as defined in Title 37, Code of Federal Regulations, §1.56(a) which became available between the filing date of the prior application and the national or PCT international filing date of this application:

U.S. Serial No.	Filing Date	Status
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I hereby appoint the following attorneys and/or agents to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith:

William E. Booth, Reg. No. 28,933
J. Peter Fasse, Reg. No. 32,983
Y. Rocky Tsao, Reg. No. 34,053
Ruffin B. Cordell, Reg. No. 33,487

Ingrid A. Beattie, Reg. No. 42,306
Janis K. Fraser, Reg. No. 34,819
Eldora L. Ellison, Reg. No. 39,967

Address all telephone calls to INGRID A. BEATTIE, PH.D., J.D. at telephone number (617) 542-5070.

Address all correspondence to INGRID A. BEATTIE, PH.D., J.D. at:

FISH & RICHARDSON P.C.
225 Franklin Street
Boston, MA 02110-2804

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patents issued thereon.

Combined Declaration and Power of Attorney
Page 2 of 2 Pages

Full Name of Inventor: Hugh S. Keeping

Inventor's Signature: _____ Date: _____
Residence Address: 10 King Philip Avenue, Bristol, RI 02809
Citizenship: Canada
Post Office Address: Same as above

Full Name of Inventor: Jonathan S. Reichner

Inventor's Signature: _____ Date: _____
Residence Address: 4 Patricia Drive, North Providence, RI 02904
Citizenship: USA
Post Office Address: Same as above

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